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Fig. 1 - in situ photographs of untreated, Ad-ßgal treated and Ad-TIMP-2 treated animals after 5 weeks;

Fig. 2 - graft showing tumor growth after treatment.

Description of the Preferred Embodiment -.

IN THE CLAIMS

Please amend the claims as follows. Claims 8, 11, 13, 15, 17, 31, and 49-51 are amended. This amendment provides a marked-up copy of the claims as amended in the Preliminary amendment filed with the application. Please note that claims 50 and 51 were not amended in the first preliminary amendment. A marked-up copy is also enclosed.

- 1. Agent for gene-therapeutic prophylaxis and therapy of tumour diseases, entailing a
- vector in the sense of a gene transfer vehicle
- enhancer/promoter
- transgene

with at least one of the components stated being aimed at the impregnation of normal tissue.

- 2. Application of the agent according to Claim 1 for genetherapeutic prophylaxis and therapy of tumour diseases, entailing a
- vector in the sense of a gene transfer vehicle
- enhancer/promoter
- transgene

with at least one of the components stated being aimed at the impregnation of normal tissue.

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- 3. Application of a gene transfer vector entailing a transgene in operative connection with an enhancer/promoter for the production of an agent for the gene-therapeutic prophylaxis and therapy of tumour diseases by administration on normal tissue.
- 4. Method for the gene-therapeutic prophylaxis and therapy of tumour diseases wherein an agent entailing a
- vector in the sense of a gene transfer vehicle
- enhancer/promoter
- transgene

with at least one of the components stated being aimed at the impregnation of normal tissue, is administered to a subject requiring a prophylactic or therapeutic tumour treatment in such a way that the vector is essentially absorbed by normal cells.

- 5. Agent according to Claim 1 with a promoter and/or enhancer regulated by transcription factors active in normal tissue.
- 6. Agent according to Claim 5 containing the CMV promoter or the SV 40 promoter or the RSV promoter, or liver-specific promoters such as the albumin promoter or lung-specific promoters or brain tissue-specific promoters or bone-specific promoters or promoters active in potential metastatic target organs or organs of the genesis of primary tumours.
- 7. Agent according to Claim 5 containing an enhancer/promoter activated by addition of an applicable substance.

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- 8. (amended) Agent according to Claim 5 which is a tetracyclindependent or a steroid hormone dependent promoter.
- 9. Agent according to Claim 1 containing transgenes for substances
- which limit the growth of the tumour
- destroy the tumour
- protect the normal tissue against tumour invasion.
- 10. Agent according to Claim 1 containing genes of metalloprotease inhibitors



11. (amended) Agent according to Claim 1 containing an antitumoral transgene coding for: TIMP-1 or TIMP-2

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12. Agent according to Claim 1 containing a protease-inhibitory transgene coding for:

TIMP-3 or TIMP-4 or PAI-1 or PAI-2.



13. (amended) Agent according to Claim 11 containing a modified transgene, the anti-tumoral effect of which has been reinforced by this modification.

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14. Agent according to Claim 13 which contains a relevant transgene C-terminal trunked TIMP-2.

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15. (amended) Agent according to Claim 1 containing a transgene of the extra-cellular matrix.



16. Agent according to Claim 15 containing at least two polypeptide chains of collagen or fibronectin or laminin or

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genes, products of which are responsible for the synthesis of non-protein components of the ECM.

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- 17. (amended) Agent according to Claim 15 containing a transgene of the extra-cellular matrix modified in such a way that it is difficult to decompose or is decomposable.
- 18. Agent according to Claim 1 containing a transgene coding for an adhesion molecule.
- 19. Agent according to Claim 18 in which the adhesion molecule in question is claudin or occludin or a cadherin or an integrin or a gene from the immunoglobulin superfamily, a selectin or a muzin.

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- 20. Agent for prophylaxis and treatment of tumour diseases containing an anti-tumoral transgene or sequences thereof which has been provided with a membrane anchor sequence.
- 21. Application of a gene transfer vector for production of an agent for prophylaxis and treatment of tumour diseases containing an anti-tumoral transgene or sequences thereof which has been provided with a membrane anchor sequence.
- 22. Method for prophylaxis and treatment of tumour diseases in which an anti-tumoral transgene or sequences thereof which has been provided with a membrane anchor sequence is transferred.
- 23. Agent according to Claim 20 containing a suicide gene or otherwise chemotherapeutically effective gene as the transgene in question

- 24. Agent according to Claim 23 in which the transgene in question is cytosin desaminase or active part sequences thereof or nitroreductase or active part sequences thereof.
- 25. Agent according to Claim 1 in which the vector is a virus.
- 26. Agent according to Claim 25 in which the virus is a first-generation adenovirus or an adeno-associated virus or a minimal adenovirus or an HSV or a lentivirus.
- 27. Agent according to Claim 26 in which the virus is a lentivirus/minimal adenovirus hybrid.
- 28. Agent according to Claim 27 in which the vector is a non-human mammal adenovirus.
- 29. Agent according to Claim 1 in which the vector is not a virus.
- 30. Agent according to Claim 29 in which the vector is a liposomal formulation or carrier proteins are used.
- 31. (amended) Agent according to Claim 25, in which the surface is modified in such a way that a specific gene transfer into the normal tissue is achieved.
- 32. Agent according to Claim 1 containing a minimal adenovirus and TIMP-2.
- 33. Agent according to Claim 1 containing a minimal adenovirus and C-terminal truncated TIMP-2.

- 34. Agent according to Claim 1 containing an AAV and TIMP-2
- 35. Agent according to Claim 1 containing a first-generation adenovirus and TIMP-2.
- 36. Agent according to Claim 1 containing a lentivirus/minimal adenovirus hybrid and TIMP-2.
- 37. Agent according to Claim 1 containing an AAV and C-terminal truncated TIMP-2.
- 38. Agent according to Claim 1 containing a minimal adenovirus and E-cadherin.
- 39. Agent according to Claim 1 containing an AAV and E-cadherin.
- 40. Agent according to Claim 1 containing a minimal adenovirus and at least two polypeptide chains of the collagen.
- 41. Agent according to Claim 1 for gene transfer into the hepatic tissue.
- 42. Agent according to Claim 1 for therapy and prophylaxis of liver metastases.
- 43. Agent according to Claim 1 for therapy of brain tumours.
- 44. Agent according to Claim 1 for therapy of lung metastases.
- 45. Agent according to Claim 1 containing the HNFAlbumin enhancer/promoter, AAV and TIMP-1.



46. Agent according to Claim 1 containing an enhancer/promoter, activated by a substance foreign to the body and containing at least two polypeptide chains of the collagen.

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- 47. Agent according to Claim 1 containing a liver-specific promoter, an AAV and a metalloprotease inhibitor.
- 48. Agent according to Claim 1 containing a liver-specific promoter, a minimal adenovirus and a metalloprotease inhibitor.
- 49. (amended) Agent according to Claim 1 containing a liverspecific promoter and a minimal adenovirus.

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- 50. (amended) Agent according to Claim 1 containing a liver-specific promoter and an AAV.
- 51. (amended) Agent according to Claim 1 containing a liverspecific promoter and a lentivirus/minimal adenovirus hybrid.

REMARKS

The above amendments were made to place the application into proper United States Patent Format.

Respectfully Submitted,

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